## APPROXIMATE THEORETICAL MODEL FOR THE FIVE ELECTRONIC STATES ( $\Omega = 5/2, 3/2, 3/2, 1/2, 1/2$ ) ARIS-ING FROM THE GROUND $3d^9$ CONFIGURATION IN NICKEL HALIDE MOLECULES AND FOR ROTATIONAL LEVELS OF THE TWO $\Omega = 1/2$ STATES IN THAT MANIFOLD

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An effective Hamiltonian for a non-rotating diatomic molecule containing only crystal-field and spin-orbit operators has been set up to describe the energies of the five spin-orbit components that arise in the ground electronic configuration of the nickel monohalides. The model assumes that bonding in the nickel halides has the approximate form Ni<sup>+</sup>X<sup>-</sup>, with an electronic  $3d^9$  configuration plus closed shells on the Ni<sup>+</sup> moiety and a closed shell configuration on the X<sup>-</sup> moiety. Least-squares fits of the observed five spin-orbit components of the three lowest electronic states in NiF and NiCl are then carried out in terms of the three crystal field parameters  $C_0, C_2, C_4$  and the spin-orbit coupling constant A. Following this, the usual effective Hamiltonian  $B(J-L-S)^2$  for a rotating diatomic molecule is used to derive expressions for the unusually large  $\Omega$ -type doubling parameter p in the two  $\Omega = 1/2$  states in the  $3d^9$  manifold. These expressions show (for certain sign conventions) that the sum of the two p values should be -2B, but that their difference can vary between -10B and +10B. The theoretical magnitudes for p are in good agreement with the two observed p values for both NiF and NiCl, but the signs are not. The experimental signs can be brought into agreement with the theoretical signs by a fairly massive change in +/- parity assignments in the NiF and NiCl literature. The last part of the talk will focus on the theoretical and experimental implications of these parity changes.